Gerald Amiel Ballena

Professional Summary

Versatile bioinformatics specialist with extensive experience in developing scalable workflows, computational biology, and machine learning. Successfully processed multi-terabyte datasets, streamlined bioinformatics pipelines to enhance reproducibility, and enabled actionable insights in environmental and public health projects.

Experience

2024 DPresent

Project Technical Specialist

University of the Philippines, College of Public Health

- O Developed scalable bioinformatics workflows for high-throughput sequence analysis (2.5 Terabytes each).
- Automated data preprocessing workflows using Snakemake, reducing manual load by 90% when data is available while also enhancing reproducibility and removing human error.
- Collaborated with cross-disciplinary teams on projects involving public health, microbiology, and science of environmental engineering.
- Enhanced data analysis pipelines, leading to actionable insights for surveillance and public health research.

Education

University of the Philippines Diliman. Graduated: July 2022 Thesis: In silico assessment of the association of pathogenicity and metal-resistance potential of *Fusarium* spp. Pre-print Link

Accomplishments: DOST ASTHRDP-Scholarship

Calculated Effective GPA (MS): 1.72

University of the Philippines Baguio.

Graduated: June 2018

Accomplishments: Advanced Placement Exam: Advanced Algebra

Philippine Science High School CAR Campus.

Graduated: March 2014

Accomplishments: Focused on STEM curriculum with a strong emphasis on research and scientific inquiry.

Skills

Technical Skills.

Programming Languages: Python (Fluent), R (Advanced), Bash (Intermediate), Perl & BioPerl (Intermediate), C++ (Fair).

Bioinformatics Tools: Kraken2, MetaPhlAn, MEGAHIT, Snakemake, Prokka, Jellyfish, BUSCO, and others.

Workflow Automation: Snakemake, Conda, YAML (Configuration Files).

Data Visualization: ggplot2, Plotly, matplotlib, QGIS.

High-Performance Computing: Slurm, Docker.

Soft Skills.

Collaboration: Proficient in leading cross-disciplinary projects and fostering effective team collaboration.

Technical Writing: Skilled in preparing technical reports and documentation using TeX tools such as TeXStudio and Overleaf.

Problem-Solving: Adept at diagnosing and optimizing bioinformatics pipelines to enhance efficiency, reproducibility, and both statistical and scientific robustness.

Certifications and Relevant Coursework

Certifications.
AI Fundamentals:
Data Literacy:
Relevant Courses.
Fundamentals: Introduction to Data Engineering
Intermediate: Bioconductor in R, RNA-Seq with Bioconductor, Differential Expression Analysis with limma (Genomic Analysis track)
Advanced: Hyperparameter Tuning in R, Designing Machine Learning Workflows in Python Nov 2024
Professional Development.
Ongoing: Ensemble Methods in Python (Bagging, Boosting, Stacking), Visualizing Geospatial Data in R.

Scripts and Workflows

Key Pipelines and Workflows: Developed and implemented scalable workflows and pipelines using Snakemake, Python, and Bash for metagenomic analysis, diversity profiling, and bioinformatics tool management. Highlights include:

- O Metagenomic Analysis Pipeline: Automated workflows for trimming, taxonomic profiling (Kraken2, Bracken), and diversity calculations (scikit-bio), ensuring high reproducibility and scalability.
- Comprehensive Binning Workflow: Designed workflows for assembly, binning (MetaWRAP, MEGAHIT), and MAG validation (CheckM2), streamlining large-scale metagenomic projects.
- Plasmid and ARG Analysis: Built pipelines for plasmid detection and antimicrobial resistance profiling using metaSPAdes, PlasmidFinder, and RGI.
- O Diversity Analysis: Developed Python-based scripts and R workflows for alpha/beta diversity metrics (Shannon, Chaoi, Bray-Curtis) and visualizations using ggplot2.

Automation and Custom Tools: Created tools and workflows for parameter optimization, repository mining, and bioinformatics tool management:

- Randomized Parameter Testing: Automated preprocessing parameter exploration for tools like Trimmomatic, Cutadapt, and fastp, enabling systematic optimization.
- Bioconda Repository Mining: Developed a Python-based scraper to extract and filter bioinformatics tools for metagenomics and AMR research.
- General Bootstrapping Workflow: Automated sampling of paired-end reads for diversity and functional analyses using seqtk.
- O Tool Management: Streamlined dependency discovery, YAML updates, and Conda-based environment

management for reproducible pipelines.

Advanced Analysis and Statistical Workflows: Designed workflows for k-mer analysis, contaminant detection, and statistical evaluations:

- K-mer Analysis: Automated frequency distribution fitting, entropy calculations, and alignment validation for metagenomic datasets (Jellyfish, MASH).
- Ocontaminant Filtering: Built pipelines for k-mer mapping and statistical testing against known contaminant databases (UniVec, PhiX, KMA).
- Visualization Pipeline: Developed R-based workflows for ridgeline and violin plots, NMDS, and heatmaps to visualize diversity and taxonomic profiles.

For a more detailed documentation of the scripts visit this link.